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


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


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Sequence concatenation is most appropriate for orthologous **sequences**. Automated identification of orthologs in **sequence database** is a challenging task both ...
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


Determining whether a **database** contains at least k genes sampled from at least ... These upper bounds to **sequence concatenation** have important implications ...
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


List **concatenation**: speed : List Concatenation « List « Python. ... `L = L + [3]` #

concatenate: slower `print L` `L.append(4)` # faster, but in-place ...

www.java2s.com › Python › List › List Concatenation - Cached - Similar -   

SQL Syntax




Mckoi SQL **Database** supports a subset of entry level ANSI SQL-92. Drops a **sequence** generator previously created with the CREATE **SEQUENCE** statement. ... INSERT INTO table SET col1 = 10, col2 = 4 + 3, col3 = **CONCAT**(col1, 'c') ...

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Efficient Algorithms for Mining Maximal Frequent **Concatenate** ...

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set of maximal frequent **concatenate sequences** in the **database**. Before introducing our method on ... **concatenate sequence** in **sequence database** S. The | - ...




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Large-Scale **Concatenation** cDNA Sequencing — Genome Research

by W Yu - 1997 - Cited by 41 - Related articles - All 11 versions




The upper size limits for a **concatenation** library that can be sequenced efficiently is not ...

Computer Search for DNA **Sequence** Similarities in the **Database** ...

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EcMLST Download **Concatenated Sequences**




Escherichia coli multilocus **sequence** typing **database**. ... In **concatenated sequence** files indexed by **sequence** types, the name of each **sequence** is : **sequence** ...

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


RibAlign: a software tool and **database** for eubacterial phylogeny ...

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


Since there is neither a comprehensive **database** for ribosomal protein **sequences** The

following 37 **sequences** were **concatenated**: RplABCDEFGHIJKLMNQRSTUUVW, ...
www.ncbi.nlm.nih.gov/pmc/articles/PMC1421441/ -   

Sequencer Pipeline

A local copy of Full implementation of Stanford Microarray **database** using ... **Sequence Concatenation** Tool. This program allows for the **concatenation** of two ...
primer.ansci.umn.edu/software1.html - Cached -   

ODBC Escape Sequences | ColdFusion Developer's Journal

ODBC escape **sequences** are a way of coding **database** functionalities that have ... but we can use an escape **sequence** that represents the **concatenation** ...
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